The mirror mechanism: recent findings and perspectives

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Mirror neurons are a specific type of visuomotor neuron that discharge both
when a monkey executes a motor act and when it observes a similar motor act
performed by another individual. In this article, we review first the basic prop-
erties of these neurons. We then describe visual features recently investigated
which indicate that, besides encoding the goal of motor acts, mirror neurons
are modulated by location in space of the observed motor acts, by the perspec-
tive from which the others’ motor acts are seen, and by the value associated
with the object on which others’ motor acts are performed. In the last part
of this article, we discuss the role of the mirror mechanism in planning actions
and in understanding the intention underlying the others’ motor acts. We also
review some human studies suggesting that motor intention in humans may
rely, as in the monkey, on the mirror mechanism.

1. Introduction

Mirror neurons were described for the first time 20 years ago in the monkey
ventral premotor area F5 [1–3], and subsequently in the monkey inferior pariet-
al area PFG [4,5]. Their discovery was preceded by a prolonged anatomical and
functional investigation of the premotor areas that enabled our group to high-
light a series of unexpected functions of these areas [6,7]. Among these
functions, were the coding of the goal of motor acts rather than movements
that form them [7,8], the responsiveness to objects in area F5 ([9] and see
[10]) and the coding of the peripersonal space in area F4 [11,12].

In area F5, besides purely motor neurons, two categories of visuomotor neur-
os were found. One, referred to as ‘canonical neurons’ [13], is responsive to the
presentation of three-dimensional objects. The other, referred to as ‘mirror neur-
os’, is responsive to the observation of motor acts performed by others [2,3].
The main property of canonical neurons is to match the shape and size of the
observed object with a specific type of prehension, whereas the main property of
mirror neurons is that of matching observation of hand and mouth motor acts
with the execution of the same or similar motor acts. This matching mechanism
enables the observing individual to achieve an automatic understanding—i.e. an
understanding without inferential processing of others’ goal-directed motor acts.

Subsequently, neurons with mirror properties were also described in corti-

cal areas outside the parietofrontal circuit, such as the mesial frontal cortex [14]
in non-human primates and the hippocampus [15] in humans. There is much
evidence that an observation/execution mechanism, demonstrated with electro-
physiological (electroencephalography (EEG), magnetoencephalography (MEG)
and transcranial magnetic stimulation (TMS)) and brain imaging (positron emission
tomography (PET) and functional magnetic resonance imaging (fMRI)) techniques,
is present in humans in areas homologous to those containing mirror neurons in the
monkey, plus the insula and the cingulate cortex (see [16–21]). This review focuses
on recent data on the functional properties of parietal and premotor mirror neurons,
including their role in intention understanding. In the last section, we discuss some
human studies dealing with this latter issue.

2. Basic properties of monkey mirror neurons

(a) Visual and motor properties

Before describing the characteristics of visual responses of mirror neurons, it is
important to stress the main properties of F5 motor neurons, including mirror...
neurons (figure 1). It is now well established that these neurons code the goal of the motor acts [28]. Evidence supporting this point is provided by neurons that discharge when the monkey grasps an object (e.g. food) with its right hand, left hand and the mouth [7]. It is clear that this type of neural behaviour cannot be explained in terms of movements. Additional evidence in favour of goal coding was provided by experiments in which the monkeys grasped the food with normal or inverted pliers (i.e. pliers that require the opening of the hand to close the pliers and therefore to grasp the object). These experiments showed that the neuron discharge correlated with the goal of the motor act, regardless of whether the hand closed or opened to achieve the goal [8]. Further studies demonstrated that the motor goal is also coded by motor neurons of the inferior parietal lobule (IPL) [5].

The main feature of the visual properties of mirror neurons in both premotor and parietal cortices is the congruence between the executed and the observed motor act [2–5]. Some mirror neurons show a strict correspondence between the effective observed and executed motor act; others show a correspondence in the goal of the observed and executed motor act, but not in the precise movements necessary to achieve the goal. Although some other specific visual characteristics have been already observed in the early studies on mirror neurons, these characteristics have been recently re-investigated and will be dealt with below.

An important aspect of mirror neuron responses that must be stressed for understanding their function is to clarify what their discharge encodes. Single neuron extracellular recordings give information about neuron output. Because the output is the same both when the monkey executes a motor act and when it observes the same motor act made by another agent, it follows that because the neuron codes the motor act during active performance, it also codes the same motor act during observation.

Further evidence that mirror neurons code the goal of others’ motor acts is provided by the experiment in which auditory stimuli typical of certain motor acts were presented to the monkey [29]. The results showed that a large number of the recorded mirror neurons responded not only to the observation of specific motor acts (e.g. breaking a peanut), but also to their sound. By contrast, other types of auditory stimuli related to different motor acts or unspecific sound-like white noise, were not effective. This finding clearly supports the notion that mirror neuron responses code the goal of motor acts performed by others.

(b) The mirror neuron circuit

It has been known, for many years, that there are many neurons in the region of the superior temporal sulcus (STS) that discharge in response to the observation of actions made by others [30]. This region, as discussed below, represents the major input to the IPL, in turn, connected to ventral premotor cortex. In the rostral part of the ventral bank of STS, some neurons were described that fired during observation of hand grasping. However, neurons in STS do not discharge during active movements. This strongly suggests that STS is the likely origin of the higher-order visual input necessary for the genesis of mirror neurons in the parietofrontal circuit.

This point has been recently investigated using fMRI complemented by neuroanatomical tracing techniques in the monkey [31]. In the fMRI experiments, video-clips of grasping actions performed by a human actor were presented. The results showed activation in three nodes (figure 2a): SPL, IPL and the arcuate region. A subsequent region of interest (ROI) analysis was carried out. For the parietal lobe, the chosen regions were the cytoarchitectonic areas PF, PFG, PG and anterior intraparietal (AIP), whereas for the STS region, they were the middle temporal area (MT) and its satellites, superior temporal polysensory middle (STPM) and two areas located in the rostral part of dorsal and ventral STS. The statistical analysis, based on the contrast between grasping action and a variety of static stimuli, showed that in the parietal lobe action-related increase in discharge was present in areas PFG and AIP. No action-specific activation was found in the other parietal areas. Action-related stimuli evoked stronger activation than static stimuli in all STS areas.

In order to elucidate which of these areas are connected with the parietal areas with mirror properties, tracers were injected in the PFG and AIP areas. The results showed a different pattern of connection of the two parietal areas with STS. Broadly speaking, area PFG was mostly connected with the upper bank of STS, and in particular with STPM, whereas AIP was mostly connected with the lower bank of STS, and in particular with the rostralmost subdivisions. Figure 2b shows the details of these connections, together with the connections of the inferior parietal areas and of STS areas to the frontal lobe. Note that the connection to AIP starts not only from STS areas, but also from a part of the cortex that is considered functionally and anatomically part of the inferotemporal cortex (see also [33]). Because inferotemporal cortex is considered to encode the meaning of the objects, these connections suggest that the mirror circuit also receives information concerning objects semantics. These data, together with previous neuroanatomical findings [33,34], indicate that there are two temporoparieto-premotor streams that process information about the actions of others. One connects the upper bank of STS/STPM with PFG and then with premotor areas F5c and F4. The other connects the lower bank of rostral STS/inferotemporal cortex with AIP and then with F5p and F5a.

3. New findings on visual properties of mirror neurons

(a) Mirror properties of corticospinal tract neurons

Kraskov et al. [35] investigated the activity of corticospinal neurons located in F5. They first identified the pyramidal tract neurons (PTNs) using antidromic stimulation. Then, they tested these neurons for their mirror properties. About half of these neurons responded when monkeys observed the experimenter grasping an object. An interesting observation was that about 25% of these PTNs showed complete suppression of discharge during grasping observation, whereas they strongly discharged during grasping execution. The authors suggested that this PTN suppression might be involved in the inhibition of the observer’s movement during observation action.

In a following study, they applied the same paradigm to PTNs recorded from primary motor cortex (F1) [36]. About half of these neurons were modulated during action observation. Of them, the majority increased their discharge during action observation (‘facilitation-type’ mirror neurons), whereas others showed reduced discharge or stopped firing.
(‘suppression-type’ mirror neurons). A comparison of the discharge of the facilitation-type neurons between grasping observation and execution showed that they discharged much less for action observation than for action execution. By comparing the properties of F1 mirror PTNs with those of F5 mirror PTNs, they observed that the visual response in F1 was much weaker than in F5. Thus, although many PTN F1 output neurons are active during action observation, their direct input to spinal circuitry may not be sufficient to produce overt muscle activity.

These data indicate that the understanding of motor goals is not only a function of F5 mirror neurons, but rather of the activation of a neural network that includes facilitatory corticospinal tract neurons.

(b) New visual properties of mirror neurons

Originally, the study of mirror neurons was focused on the match between visual and motor responses. This matching is, indeed, fundamental for the hypothesis that these neurons play a role in understanding the goal of motor acts. However, already in the early studies, some subcategories of mirror neurons, based on their specific visual properties, were identified. These subcategories included neurons showing selectivity for specific space positions, right or left hand, and for specific directions [3]. These previous observations have been recently extended, as will be described below.

(i) Space-sensitive mirror neurons

One of the issues addressed in these new studies is that of the possible influence on mirror neuron discharge of the spatial location of the observed actions. Caggiano et al. [37] performed a study in which they tested mirror neurons recorded from area F5, while the experimenter grasped objects in either the space around the monkey (peripersonal space) or in the space outside its reach (extrapersonal space; figure 3a). All neurons were also tested during active grasping by the monkey.

The results showed that the visual response of half of the recorded mirror neurons was influenced by the location in space of the observed motor acts, whereas the others did not show a differential modulation. Of those sensitive to action location in space, about half preferred the monkey’s peripersonal space, the other half the extrapersonal space. Examples of the two types of space-selective neurons and one unselective neuron are shown in figure 3b.

Subsequently, the authors investigated whether space-selective neurons encode space in a ‘metric’ or in an ‘operational’ format. Metric format means that there is a fixed boundary between the two spaces, whereas operational format means that the space is dynamic and depends on the monkey’s possibility to reach the objects. To study this issue, monkeys were tested with motor acts performed by the experimenter in the peripersonal and extrapersonal space. When neurons discharging differentially in the two space sectors were identified, a frontal transparent barrier was introduced that prevented the monkey from reaching for objects close to his body. About half of the tested space-selective mirror neurons were found to be influenced by the experimental manipulation (operational mirror neurons). More specifically, after closure of the transparent barrier, these neurons, when selective for the extrapersonal space, also started to respond in the peripersonal space, whereas when selective for the peripersonal space, they ceased to respond (figure 3c). In other words, the introduction of the barrier blocked the monkey’s possibility to act in its reaching space, thus shrinking its space representation. As a consequence, an observed action performed in its peripersonal space was now located in the new extrapersonal space. The other half of neurons code space in a metric way (Cartesian mirror neurons) and were not influenced by the introduction of the panel.

Figure 1. Lateral view of the monkey brain showing the subdivisions of the agranular frontal and posterior parietal cortices. The intraparietal and arcuate sulci have been opened to show the areas buried inside them. Agranular frontal areas have been labelled according to Matelli et al. [22,23]. Note that area F5 is formed by three further subdivisions: F5c, F5p and F5a [24]. Posterior parietal areas are defined according to Pandya and Seltzer and Gregoriou et al. [25,26]. The areas buried inside the intraparietal sulcus are defined according to functional criteria (for references, see [27]). AI, inferior arcuate sulcus; AIP, anterior intraparietal area; AS, superior arcuate sulcus; C, central sulcus; DLPF, dorsolateral prefrontal cortex; FEF, frontal eye field; IO, inferior occipital sulcus; L, lateral fissure; LIP, lateral intraparietal area; Lu, lunate sulcus; MIP, medial intraparietal area; P, principal sulcus; ST, superior temporal sulcus; VIP, ventral intraparietal area; VLPF, ventrolateral prefrontal cortex.
Figure 2. (a) Overview of MR brain activations (recorded with 3T fMRI) during the observation of grasping acts. Leftmost image: lateral view of reconstructed left hemisphere indicating the six different antero-posterior levels at which coronal slices shown on the right have been taken. Right: statistical parametric maps activation for the contrast: hand action versus static control. The data are from a single monkey, overlaid onto its coronal anatomical sections. Numbers on each slice indicate y-coordinate (antero-posterior from interaural plane). as, arcuate sulcus; ips, intraparietal sulcus; sts, superior temporal sulcus. (b) Temporo-parieto-premotor grasping observation pathways in the monkey brain. Flattened representation of STS, IPS/IPL and IAS with ROIs indicated. Arrows and areas coloured in red and blue indicate, respectively, the STPm–PFG–F5c pathway and the LB2–AIP–F5a/p pathway. For the definition of areas PF, PFG, PG and Opt, see figure 1. Areas 45a and 45b are defined according to Gerbella et al. (32). AIP, anterior intraparietal area; FEF, frontal eye fields; F5c, F5 convexity; F5p, F5 (bank) posterior; F5a, F5 (bank) anterior; FST, fundus of the STS; IAS, inferior arcuate sulcus; LIPa, anterior part of the lateral intraparietal area; MT/V5, middle temporal area; MSTd, middle superior temporal area, dorsal part; MSTv, middle superior temporal area, ventral part.
A possible functional role of space-sensitive mirror neurons is that they may set which is the most appropriate behavioural reaction according to the location of the observed action in space. Peripersonal space suggests a possible immediate interaction with the action agent, whereas extrapersonal space implies a more complex behavioural pattern in order to interact with the acting individual. This interaction could be cooperative or competitive.

An interesting issue is which mechanism it is that may determine the spatial properties of these neurons. The most likely candidate is area F4. Neurons in this area respond to tactile, visual and more rarely, auditory stimuli [11,12,38,39]. Their visual receptive fields (RFs) are located around the tactile ones and extend into the monkey peripersonal space. Most of these RFs are ‘operational’, in the sense that their depth extent is modulated by the speed of an approaching stimulus [12].

Figure 3. Space-selective mirror neurons. (a) Schematic view of the visual conditions of the experimental paradigm: the monkey observes an experimenter executing a grasping act in the peripersonal (left) and extrapersonal (right) space of the monkey. (b) Examples of the responses of three mirror neurons during observation of grasping acts executed in the monkey’s peri- and extrapersonal space and during monkey execution. Each panel shows the raster plot (top) and the cumulative histogram (bottom) of the neuron responses. Raster plots and histograms are aligned with the time of contact of the experimenter’s or monkey hand with the object. Neuron 1 responds more strongly when the observed grasping is performed in the extrapersonal space, while neuron 2 presents a stronger discharge during observation of grasping performed in the peripersonal space. Neuron 3 does not show any space-selective visual response. All neurons discharge during grasping execution. (c) Operational encoding of the monkey peri- and extrapersonal space. The top part shows the experimental conditions: the experimenter grasps an object in the extrapersonal (left) or in the peripersonal space without (centre) or with (right) a frontal panel impairing the monkey’s reach into its peripersonal space. Note that in this latter condition, the object (and the act performed by the experimenter) is metrically in the monkey’s peripersonal space, but operationally outside it. The lower panels show the visual responses of a mirror neuron in the three conditions. The vertical lines mark the time of contact between the experimenter’s hand and the object. Before closure with the frontal panel, the neuron was activated only during observation of grasping in the monkey’s extrapersonal space. However, after closure, the neuron also responded to observation of grasping performed close to the monkey’s body.
Because information coming from F4 reaches the adjacent area F5 [40], one might postulate that this input gates the visual responses of ‘operational’ mirror neurons.

(ii) View-dependent mirror neurons

Recently, it was investigated whether mirror neurons can also provide information concerning the perspective from which the motor acts of others are observed [41]. Preliminary to this study was an investigation aimed to establish the capacity of mirror neurons to respond not only to actions performed by an experimenter, as done in earlier studies, but also to video-clips representing hand actions. The results showed that, although the responses to video-clips were typically weaker than to live actions, nevertheless they were consistent and strong enough to be studied. Only a minority of F5 mirror neurons did not respond to video-clip presentation. These results were fundamental for studying the responsiveness of neurons to different perspectives because it allowed a standardized presentation of the stimuli.

After this preliminary investigation, mirror neurons of monkey area F5 were tested while presenting movies showing grasping motor acts from different visual perspectives (figure 4a). The three tested perspectives were subjective (0°), side (90°) and frontal (180°). The results showed that the majority of the tested mirror neurons (74%) were view-dependent, with responses tuned to one or, more frequently, to two specific points of view. Among the neurons specific for only one view, there was a slight, although not significant, preference for the subjective view. Finally, a minority of the tested mirror neurons (26%) exhibited view-independent responses. Examples of the four categories of neurons are shown in figure 4b.

A classical view in visual physiology is that view-independent neurons are generated by convergence of several lower-order neurons with view-dependent properties. In particular, this was postulated for STS neurons responding to biological stimuli, such as different head perspectives [42]. Thus, one may suggest that view-dependent mirror neurons in F5 represent an intermediate step leading to the formation of the view-independent ones. However, one must remember that mirror neurons are motor neurons, and their output target is always the same, regardless of what is the input triggering it. The message conveyed to other centres does not depend, therefore, on how a given neuron was triggered and whether the stimulus was subjective, side or frontal. Thus, both view-independent and view-dependent mirror neurons encode action goals, irrespective of the details of the observed motor acts.

What may be therefore the functional meaning of the view-dependent mirror neurons? An interesting possibility is that view-dependent mirror neurons play a role in the perception of the visual perspective of the observed actions. There is anatomical evidence that, in addition to feed-forward projections, the nodal areas of the mirror circuit plus STS are also connected by backward projections [33,34]. Hence, signals from view-dependent mirror neurons might contribute to action perception by modulating the activity of visual representations coded by STS neurons, reinforcing the processing of visual patterns that are associated with the different views of a grasping act. This hypothesis suggests that the understanding of action goal is carried out by mirror neurons, whereas the perception of specific details of the observed action is provided by higher-order visual neurons, following facilitation coming from mirror neurons (for a similar view in visual perception, see Hochstein & Ahissar [43]).

(iii) Mirror neurons sensitive to the value of an observed action

Objects grasped by an agent have a value not only for the individual performing that act, but also for the individual...
observing it. A recent study addressed the problem of whether F5 mirror neurons are involved in representing the value of the grasped object [44]. Three experiments were carried out.

In the first experiment, the responses of mirror neurons to the observation of grasping food were compared with those to the observation of grasping objects devoid of any value for the monkey. The results showed that about 70% of the studied mirror neurons were influenced by the type of the observed grasped object. More specifically, the large majority of these neurons were more strongly activated when grasping was directed towards food than when it was directed towards the object devoid of value.

These findings were confirmed in a second experiment, where the responses of mirror neurons were studied in response to the observation of grasping motor acts on objects associated with reward or on similar objects, but not rewarded. The data showed that about half of the tested mirror neurons exhibited stronger visual responses when the observed motor acts were performed on rewarded objects. Only a small percentage of neurons showed a stronger activity for non-rewarded objects. Finally, about 40% of neurons were not modulated by the presence or absence of reward associated with grasped objects.

In the third experiment, it was tested whether mirror neurons reflect differences in subjective value in a continuous rather than a categorical manner. In order to address this issue, the experiment was carried out by presenting to the monkey a series of blocks of grasping trials in which the experimenter grasped a non-food object. On a given block, the motor act directed to the object could be associated with one out of three types of consequences for the observing monkey: (i) the delivery of highly palatable food, (ii) the delivery of less palatable food or (iii) no reward at all. The results showed that about 50% of the tested neurons exhibited a significant effect of the stimulus subjective value. Among these value-sensitive mirror neurons, a large number (35%) showed selectivity for the favourite reward condition, whereas about half did not show any preference for the type of reward. Finally, a few neurons (about 15%) discharged stronger in response to the less favourite reward or even to the absence of a reward.

At first glance, this influence of the subjective value on mirror neuron responses may sound surprising. The traditional view on mirror neuron activity was that they were activated by the type of motor act, but not by the semantics of the object on which the motor act was performed. However, there is evidence from both fMRI and connectivity studies that one of the nodes of the mirror system (parietal area AIP) receives information not only from the lower bank of STS, but also from the inferotemporal lobe. Thus, in the case of grasping, when the observed grasped object is food, the neurons reflect differences in subjective value in a continuous rather than a categorical manner. The results showed that about 50% of the tested neurons exhibited a significant effect of the stimulus subjective value. Among these value-sensitive mirror neurons, a large number (35%) showed selectivity for the favourite reward condition, whereas about half did not show any preference for the type of reward. Finally, a few neurons (about 15%) discharged stronger in response to the less favourite reward or even to the absence of a reward.

Another important distinction is that between motor goal and motor intention. We define as motor goal the outcome of a motor act, whereas we define as motor intention the final outcome of an action that leads to positive or negative reinforcement. Although this distinction is somehow implicit in the definition of motor act and action, it is important to stress it because, frequently, the two terms are confused with each other.

4. Motor intention and mirror neurons

It is important at the outset of this section to clarify some terminological aspects. We consider ‘movement’, ‘motor act’ and ‘action’ as terms describing different aspects of motor behaviour. We define as movement sensu stricto a simple joint displacement or orofacial twitches. For example, a finger flexion elicited by electrical or magnetic stimulation represents a typical movement. Motor act, we define as a series of movements organized in such a way as to achieve a specific motor goal. For instance, shaping of the hand in such a way as to grasp an object represents a typical motor act. Finally, we define an action as a sequence of fluently linked motor acts that, as its final outcome, determines the achievement of a behavioural goal. For example, reaching, grasping and bringing a piece of food to the mouth for eating represent a typical action.

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(a) Coding of motor sequences in the parietal and premotor cortex

The same motor act may be part of different actions having different final behavioural goals. In order to assess the neural basis of this motor organization, single neurons were recorded from the IPL and tested in different actions [4].

Neurons discharging during the execution of grasping motor acts were tested in two main conditions. In the first, the monkey had to reach and grasp a piece of food and bring it to the mouth; in the second, the monkey had to reach and grasp an object and place it into a container (figure 5a, left). In the first condition, the monkey was allowed to eat the grasped food; in the second condition, it was rewarded with a piece of food given to it by the experimenter.

The results showed that the discharge of the large majority of the recorded neurons was modulated depending on the action in which the grasping act was embedded. Examples are shown in figure 5b. Unit 67 discharged during grasping, when it was followed by bringing the food to the mouth. By contrast, the neuron was virtually silent when grasping was followed by placing the object into the container. Unit 161 is an example of a neuron discharging preferentially during grasping-to-place. Finally, unit 154 discharged in the same way in the two conditions.

In total, about two-thirds of neurons discharged preferentially when grasping was embedded into a specific action. Of them, the great majority (73%) preferred grasping for eating.
Figure 5. (a) Motor task. The monkey, starting with its hand from a fixed position (left), reaches and grasps a piece of food (or an object), then it brings the food to the mouth and eats it (grasp-to-eat condition I) or places it (or the object) into a container (grasp-to-place condition) located near the mouth (II) or near the target (III). Visual task. The experimenter, starting with his hand from a fixed position (left), reaches and grasps a piece of food or an object (right), then he brings the food to the mouth and eats it (grasp-to-eat condition I) or places it (or the object) into a container located near the target (grasp-to-place condition II).

(b) Examples of the discharge of three IPL neurons during the motor task. Neuron 67 is selective for grasping to eat, neuron 161 shows the opposite behaviour, while the response of neuron 158 is not affected by the action goal.

(c) Examples of the discharge of three IPL neurons during the visual task. Neuron 87 discharges stronger during observation of grasping to eat, neuron 39, on the contrary, during observation of grasping to place, while neuron 80 discharges equally well in both conditions. In both (b) and (c), rasters and histograms are aligned (vertical bar) with the moment when the monkey or the experimenter, respectively, touched the food/object.

(d) Congruence between the visual and the motor preference

(i) Example of a neuron discharging stronger during grasping for eating than during grasping for placing, both when the action is executed and when it is observed. Conventions as in (b) and (c).

(ii) Population-averaged responses, showing the same pattern of differential activity between the preferred (white bar) and not preferred (black bar) action during motor and visual tasks. Depending on the neuron, the preferred action could be grasp-to-eat or grasp-to-place.
In order to control that the differential discharge of neurons during the same motor act performed in the two conditions was not due to a mere difference in the stimuli to be grasped, monkeys were trained to grasp the same piece of food in both conditions. The results showed that neuron selectivity remained the same, not depending on the stimulus used.

Human studies have shown that the first component of an action is influenced by subsequent components of that action [49–53]. In other words, programming of a motor act takes into account the constraints posed by the target of the subsequent motor act. This point was tested in the two conditions of the experiment described above, measuring the kinematics of the reach-to-grasp motor act. It was found that this motor act, when followed by arm flexion (bringing the food to the mouth) was faster than the same motor act when followed by arm abduction (placing the food into the container). A third experimental condition was therefore introduced to render the reaching-to-grasp movement for placing kinematically similar to that for eating. In this new condition, the monkey had to grasp a piece of food and place it into a container located near the monkey mouth, so that the animal was required, as in the bringing to the mouth condition, to flex the arm. The results showed that the arm velocity in the new placing condition was faster both than in the original placing condition as well as than in the grasping-to-eat condition. These findings indicate that the differential discharge during grasping in the two actions did not depend on kinematic parameters.

In a subsequent study, neurons were recorded from both F5 and the IPL (area PFG) in order to compare the activity of F5 and PFG in motor sequence organization [54]. The same conditions as in the study described above were used. The results confirmed that a large number of F5 neurons code grasping according to the goal of the action in which it is embedded. The same effect was found also in PFG. A comparison of the neuronal properties of the two areas showed that there is a larger percentage of grasping neurons in area PFG that are modulated by the action in which they are embedded and that they are more strongly influenced by the action goal than neurons in F5.

It may sound surprising that most IPL neurons discharge differently for the same motor act depending on the final action goal. This arrangement might seem highly inefficient, because a large number of neurons with similar properties are required for executing different types of actions. There is, however, another aspect of motor organization that must be taken into consideration: the fluidity with which one motor act follows another. This fluidity is necessary, because action execution should occur without any gap. The results of the above-described studies suggest that neurons encoding specific motor acts within an action form pre-wired intentional chains, in which a neuron encoding a motor act is facilitated by the neuron encoding the previous one.

This type of action organization raises two questions. The first one is whether more complex actions present the same type of chain organization. The second one is how this high-level, goal-centred organization is translated into actual movement organization, in which kinematic parameters must be taken into consideration.

In order to answer the first question, an experiment was carried out in which monkeys were trained to perform complex action sequences [55]. Specifically, the monkey had to grasp the lid of a container and subsequently grasp the food or the object inside it in order to eat the food (first condition, eating) or to place the object into another container (second condition, placing). Neurons were recorded from both area PFG and area F5. Confirminng the previous studies, a large number of PFG and F5 neurons displayed action-goal selectivity during the second grasping act, when the vision of the target allowed the monkey to perform the last part of the action (bringing to the mouth or placing). Notably, a small percentage of PFG neurons, but not neurons in F5, reflected the final goal from the early phase of action unfolding, when only memory-driven information was available. On the other hand, when monkeys could not see the target before action onset, neurons lost their early selectivity. These findings suggest that a higher-order chain organization does exist in PFG, but it is likely that other cortical areas, such as prefrontal cortex, play a crucial role in this organization.

As far as the second question is concerned, we have no empirical data to show how goal-related organization is translated into a kinematic-related organization. We know, however, as it will be described later, that a chain organization similar to that described at the neuronal level in the monkey also exists in humans and is reflected in the movement kinematics [56].

(b) Coding of observed motor sequences in the parietal and premotor cortex

As already mentioned, in the IPL, there are a considerable percentage of mirror neurons discharging both during grasping observation and grasping execution. In the same study in which the motor properties of IPL neurons were investigated during execution of action sequences, grasping mirror neurons were tested in a visual task in which the experimenter performed, in front of the monkey, the same actions that the monkey performed in the motor task, for example, grasping to eat and grasping to place [4]. The monkey had simply to observe the performed actions (figure 5c, right).

The results showed that while the discharge of some neurons was not influenced by the motor act following the observed grasping, the majority (75%) of mirror neurons discharged differently according to whether the grasping made by the experimenter was followed by bringing to the mouth or by placing. Examples are shown in figure 5c. Neurons responding to the observation of grasping for eating were the most commonly encountered.

In a subsequent experiment, grasping mirror neurons were recorded from area F5 as well as from area PFG, with the same experimental paradigm used in the study described above [54]. It was found that, as in PFG, also in F5, most grasping mirror neurons discharged differently during observation of grasping when this act was embedded into different actions. In both F5 and PFG, the large majority of neurons discharged more strongly during the eating condition than during the placing condition, but the prevalence of ‘eating-related’ mirror neurons was larger (90% versus 80%) in F5 than in PFG.

In a group of F5 and PFG mirror neurons, visual and motor selectivity for eating or placing was compared. The results showed that the great majority of the tested neurons showed the same specificity during grasping observation and grasping execution (figure 5d).

The most accepted interpretation of the role of mirror neurons is that they allow one to understand the goal of an
observed motor act (see for review [57]). The above-described findings point to a new mechanism, based on mirror neurons, that allows one to encode differently two identical observed grasping motor acts according to the action in which these acts are embedded. This ability, based on context or on the past history (e.g. blocked design), enables the individual to understand others’ intentions. More specifically, the selection of a specific population of grasping neurons activates the same motor chain that the observers would activate when performing the observed action themselves. As discussed before concerning action execution, the activation of a given chain corresponds with a specific intention. Thus, the observation of a motor act, which triggers a given chain, would provide the observer with the capacity to understand the agent’s specific intention. This mechanism allows an automatic, not inferential understanding of the motor intentions of others.

5. Coding motor intention in humans

In comparison with the enormous amount of studies devoted to action observation (see for review [16–21]), only a limited number of papers addressed the issue of understanding motor intention in humans. An early attempt to investigate the relation between intention coding and the mirror mechanism was done by Iacoboni et al. [58]. This fMRI study was formed by three experimental conditions. In the first, referred to as ‘context’, the subjects were instructed to observe a scene composed of objects arranged either as for the beginning of a breakfast or as for the end of it. In the second, referred to as ‘action’, the subjects had to observe a hand grasping a mug on an empty background. In the third condition, referred to as ‘intention’, the subjects had to observe the same hand motor act embedded in one of the two contexts of the ‘context’ condition. The context suggested the agent’s intention, and specifically the scene representing breakfast beginning suggested that the intention of the agent was to grasp the mug for drinking, whereas the scene representing breakfast end suggested that the intention of the agent was to grasp it for cleaning the table. The results showed that areas belonging to the cortical mirror system were activated in both ‘action’ and ‘intention’ conditions. However, the contrast between ‘intention’ and ‘action’ conditions showed activation in the right caudal inferior frontal gyrus.

The conclusion that the right hemisphere is involved in understanding the intention of others has been also reached by Ortigue et al. [59] using high-density EEG study. They investigated the temporal dynamics of brain activations produced by the observation of motor acts of subjects instructed to understand the agent’s intention. The results revealed that after an initial bilateral activation of occipital and temporo-parietal areas, there was a selective activation of the right temporo-parietal region. This late activation was interpreted by the authors as related to the understanding of intentions of others.

In a study prompted by the findings of Fogassi et al. [4] in the monkey (see above), Cattaneo et al. [56] asked typically developing children to grasp a piece of food for eating it or to grasp a piece of paper for placing it into a container. In a subsequent experimental condition, the same children had to observe an experimenter performing the same actions. In both conditions, the electromyography activity of a muscle involved in mouth opening (mylohyoid muscle, MH) was recorded. It was found that during execution there was an increase of MH activity already during the phase in which the hand was approaching the object, about 800 ms before actual grasping. A precocious activation of MH was also found during the observation of the eating action. By contrast, no MH activity increase was found during execution and observation of placing. These findings suggest that humans are endowed with a chain organization similar to that of the monkey (see above), allowing, on the one hand, the execution of fluent motor actions, and on the other hand, most interestingly, the understanding of motor intentions of others.

The studies discussed above indicate that the mirror mechanism might explain some instances of intention understanding. However, intention understanding encompasses different levels, ranging from motor intention comprehension to intention understanding based on propositional attitudes such as beliefs or desires. Thus, other mechanisms, beside that based on the mirror mechanism, are likely to be involved.

Recent fMRI experiments support these theoretical considerations. Brass et al. [60] asked subjects to observe several types of unusual actions in plausible and implausible contexts. An activation of the mirror network was present in both context conditions. However, when comparing actions performed in an implausible context versus actions performed in a plausible context, there were additional activations of the STS region and of the anterior mesial frontal cortex (see also [61] for similar observations). The activated areas are considered to be part of the so-called mentalizing network [62], that is that network that becomes active when an observer predicts someone else’s behaviour based on his/her belief state.

We mentioned that there are different levels of intention understanding. They should correspond to different mechanisms. As discussed in this section, understanding action intention relies on the mirror mechanism and the motor chain organization of the cortical motor system. By contrast, understanding the reasons underlying a given action appears to be localized in areas, such as the anterior mesial frontal cortex and the right temporo-parietal junction, which do not seem to have mirror properties. Several attempts have been made to integrate these two kinds of intention understanding (see [62]). However, it is important to stress that there are currently no neurophysiological data that can account for the mechanisms underlying the ‘mentaling network’.

6. Conclusion

The mirror mechanism has raised great interest over the past two decades, because it allows one to unify action production and action observation. This motor-based action and intention understanding appears to represent the primary way for inter-individual interactions. This is corroborated by its presence not only in humans and monkeys, but also in evolutionarily distant vertebrate classes, such as birds [63,64]. It would be of great interest to learn whether the mirror mechanism also exists in other mammalian species. Its demonstration in rodents would be particularly interesting. Besides its theoretical interest, this would allow the characterization of the mirror mechanism at a neurochemical level.

Another extremely interesting issue to be addressed is the relationship between the basic mirror circuit and other circuits that, in the monkey, can exert a top-down control. The data from Buccino et al. [65] in humans suggest that during imitation
learning, the prefrontal lobe (area [46]) is playing a fundamental role in the reorganization of the basic information about the motor acts of others captured by the mirror system. A similar kind of ability could be studied in the monkey to understand in depth how the interaction between prefrontal lobe and the parieto-premotor mirror circuit takes place. In a similar vein, it will be of great interest to assess in more controlled conditions the findings of Mukamel et al. [15] in humans showing the activation of hippocampal cells during action observation, and in particular whether this hippocampal activation is related to episodic memory.

Finally, the link between the mirror mechanism and mentalizing will be achieved only if it becomes possible to record single neurons from awake human beings during such processes. Currently, this technology has been used only in some laboratories in epileptic patients, exploiting special electrodes placed intracortically in deep mesial structures [15,66]. It is likely that in the near future modifications of classical macroelectrodes for intracortical EEG recordings will also allow the recording of single neurons from other neural structures, thus enabling the investigation of neuronal responses during cognitive tasks, in particular those requiring reasoning about the behaviour of others.

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